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1

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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.	
10/828,557	04/21/2004	Douglas G. Storey	028722-381	5430	
41790 75	90 7590 07/13/2006		EXAMINER		
BUCHANAN, INGERSOLL & ROONEY LLP			ZEMAN, R	ZEMAN, ROBERT A	
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ALEXANDRIA, VA 22313-1404			ART UNIT	PAPER NUMBER	
			1645		
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Please find below and/or attached an Office communication concerning this application or proceeding.

		Application No.	Applicant(s)			
		10/828,557	STOREY ET AL.			
	Office Action Summary	Examiner	Art Unit			
		Robert A. Zeman	1645			
	The MAILING DATE of this communication appears on the cover sheet with the correspondence address Period for Reply					
WHIC - Exte after - If NC - Failu Any	ORTENED STATUTORY PERIOD FOR REPLY CHEVER IS LONGER, FROM THE MAILING DANS INSTRUCTION OF THE MAILING DANS OF THE MAILING DANS OF THE MONTHS FROM THE MAILING DANS OF THE MONTHS FROM THE MAILING DANS OF THE MONTHS FROM THE MONTHS FROM THE MONTHS OF THE MO	ATE OF THIS COMMUNICATION 36(a). In no event, however, may a reply be tim vill apply and will expire SIX (6) MONTHS from cause the application to become ABANDONE	N. nely filed the mailing date of this communication. D (35 U.S.C. § 133).			
Status						
1)⊠ 2a)⊠ 3)⊟	Responsive to communication(s) filed on <u>15 Ma</u> This action is FINAL . 2b) This Since this application is in condition for allowant closed in accordance with the practice under <i>E</i>	action is non-final. nce except for formal matters, pro				
Disposit	ion of Claims					
5)□ 6)⊠ 7)□	Claim(s) 1-18 is/are pending in the application. 4a) Of the above claim(s) 4-7 and 9-18 is/are w Claim(s) is/are allowed. Claim(s) 1-3 and 8 is/are rejected. Claim(s) is/are objected to. Claim(s) are subject to restriction and/or	ithdrawn from consideration.				
Applicat	ion Papers					
10)⊠	The specification is objected to by the Examine The drawing(s) filed on <u>18 April 2006</u> is/are: a) Applicant may not request that any objection to the Replacement drawing sheet(s) including the correction The oath or declaration is objected to by the Ex	☑ accepted or b)☐ objected to define accepted or b)☐ objected to define acceptance. See ion is required if the drawing(s) is object.	e 37 CFR 1.85(a). lected to. See 37 CFR 1.121(d).			
Priority (under 35 U.S.C. § 119					
 12) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f). a) All b) Some * c) None of: 1. Certified copies of the priority documents have been received. 2. Certified copies of the priority documents have been received in Application No 3. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)). * See the attached detailed Office action for a list of the certified copies not received. 						
Attachmen	• •	" □	(DTO 442)			
2) Notice (3) Information	ce of References Cited (PTO-892) ce of Draftsperson's Patent Drawing Review (PTO-948) mation Disclosure Statement(s) (PTO-1449 or PTO/SB/08) er No(s)/Mail Date	4) Interview Summary Paper No(s)/Mail Da 5) Notice of Informal F 6) Other:				

The amendment and response filed on 3-15-2006 are acknowledged. Claim 1 has been amended. Claims 1-18 are pending. Claims 4-7 and 9-18 remain withdrawn from consideration as being drawn to non-elected inventions. Claims 1-3 and 8 are currently under examination.

Drawings

The drawings were received on 4-18-2006. These drawings are accepted.

Claim Objections Withdrawn

The objection to the specification based on an improper incorporation by reference is withdrawn in light of the amendment thereto.

Claim Objections Maintained

The objection to Claims 1, 2 and 8 for reciting material drawn to non-elected inventions is maintained. The elected invention (Group I of the restriction) is drawn to methods of preventing biofilm formation utilizing antibodies to gacS. Claims 1, 2 and 8 encompass other means (other than anti-gacS antibodies) to prevent biofilm formation.

Claim Rejections Maintained

35 USC § 112

The following is a quotation of the first paragraph of 35 U.S.C. 112:

The specification shall contain a written description of the invention, and of the manner and process of making and using it, in such full, clear, concise, and exact terms as to enable any person skilled in the art to which it

pertains, or with which it is most nearly connected, to make and use the same and shall set forth the best mode contemplated by the inventor of carrying out his invention.

The rejection of claims 1-3 and 8 under 35 U.S.C. 112, first paragraph, as failing to comply with the written description requirement is maintained for reasons of record. The claim(s) contains subject matter which was not described in the specification in such a way as to reasonably convey to one skilled in the relevant art that the inventor(s), at the time the application was filed, had possession of the claimed invention.

Applicant argues:

- 1. GacS is a known protein.
- 2. According to the USPTO guidelines if the structural characteristics of a protein are known, antibodies to such a protein a sufficiently known and described.

Applicant's arguments have been fully considered and deemed non-persuasive.

Contrary to Applicant's assertion the sequence of gacS is not known protein sequence associated with it. As evidenced by the attached GenBank sequence listings, gacS (also known as LemA and ApdA) have multiple sequences associated with it. Consequently, said protein has not been fully characterized. The courts have recently decided in Randolph J. Noelle v Seth Lederman, Leonard Chess and Michael J. Yellin (CAFC, 02-1187, 1/20/2004) that a patentee of a biotechnological invention cannot necessarily claim a genus after only describing a limited number of species because there may be unpredictability in the results obtained from species other than those specifically enumerated. See Enzo Biochem II, 323 F.3d at 965; Regents, 119 F.3d at 1568. Therefore, based on our past precedent, as long as an applicant has disclosed a

"fully characterized antigen," either by its structure, formula, chemical name, or physical properties, or by depositing the protein in a public depository, the applicant can then claim an antibody by its binding affinity to that described antigen. Noelle did not provide sufficient support for the claims to the human CD40CR antibody in his '480 application because Noelle failed to disclose the structural elements of human CD40CR antibody or antigen in his earlier '799 application. Noelle argues that because antibodies are defined by their binding affinity to antigens, not their physical structure, he sufficiently described human CD40CR antibody by stating that it binds to human CD40CR antigen. Noelle cites Enzo Biochem II for this proposition. This argument fails, however, because Noelle did not sufficiently describe the human CD40CR antigen at the time of the filing of the '799 patent application. In fact, Noelle only described the mouse antigen when he claimed the mouse, human, and genus forms of CD40CR antibodies by citing to the ATCC number of the hybridoma secreting the mouse CD40CR antibody. If Noelle had sufficiently described the human form of CD40CR antigen, he could have claimed its antibody by simply stating its binding affinity for the "fully characterized" antigen. Noelle did not describe human CD40CR antigen. Therefore, Noelle attempted to define an unknown by its binding affinity to another unknown. As a result, Noelle's claims to human forms of CD40CR antibody found in his '480 application cannot gain the benefit of the earlier filing date of his '799 patent application.

As demonstrated above, Applicant has failed to "fully characterize" the gacS protein to which the claimed antibody binds. Consequently, since Applicant has not fully characterized the antigen to which the claimed antibodies bind, the written description requirements under 35 U.S.C 112, first paragraph have not been met.

As outlined previously, the rejected claims are drawn to the use of antibodies specific for gacS to prevent biofilm formation wherein said antibodies inhibit the gacA/gacS regulatory system. Hence, the claims are drawn to a vast genus of genus of antibodies, the members of which recognize the gacS protein wherein said antibodies can prevent biofilm formation. To fulfill the written description requirements set forth under 35 USC § 112, first paragraph, the specification must describe at least a substantial number of the members of the claimed genus, or alternatively describe a representative member of the claimed genus, which shares a particularly defining feature common to at least a substantial number of the members of the claimed genus, which would enable the skilled artisan to immediately recognize and distinguish its members from others, so as to reasonably convey to the skilled artisan that Applicant has possession the claimed invention. To adequately describe the genus of antibodies that bind to the gacS protein, one must describe not just the antigenic determinants (immunoepitopes) of said protein.

The specification does not describe with any degree of specificity a single member of the genus of immunoepitopes of gacS to which the members of the claimed genus of antibodies must bind, wherein said antibodies can effectively prevent biofilm formation such that the specification might reasonably convey to the skilled artisan that Applicant had possession of the claimed invention at the time the application was filed.

Moreover, the specification does not disclose distinguishing and identifying features of a representative number of members of the genus of antibodies to which the claims are drawn, such as a correlation between the structure of the immunoepitope its recited function (to prevent biofilm formation), so that the skilled artisan could immediately envision, or recognize at least a substantial number of members of the claimed genus of antibodies. Additionally, the

specification fails to disclose which amino acid residues are essential to the function of the immunoepitope or which amino acids might be replaced so that the resultant immunoepitope retains the activity of its parent, or by which other amino acids the essential amino acids might be replaced so that the resultant immunoepitope retains the activity of its parent. Therefore, since the specification fails to adequately describe at least a substantial number of members of the genus of immunoepitopes on which the claims are based; the specification fails to adequately describe at least a substantial number of members of the claimed genus of antibodies that bind to the gacS protein and have prophylactic efficacy against biofilm formation.

MPEP § 2163.02 states, "[a]n objective standard for determining compliance with the written description requirement is, 'does the description clearly allow persons of ordinary skill in the art to recognize that he or she invented what is claimed'". The courts have decided:

The purpose of the "written description" requirement is broader than to merely explain how to "make and use"; the applicant must convey with reasonable clarity to those skilled in the art that, as of the filing date sought, he or she was in possession of the invention. The invention is, for purposes of the "written description" inquiry, whatever is now claimed.

See Vas-Cath, Inc. v. Mahurkar, 935 F.2d 1555, 1563-64, 19 USPQ2d 1111, 1117 (Federal Circuit, 1991). Furthermore, the written description provision of 35 USC § 112 is severable from its enablement provision; and adequate written description requires more than a mere statement that it is part of the invention and reference to a potential method for isolating it. See Fiers v. Revel, 25 USPQ2d 1601, 1606 (CAFC 1993) and Amgen Inc. V. Chugai Pharmaceutical Co. Ltd., 18 USPQ2d 1016.

The Guidelines for Examination of Patent Applications Under the 35 U.S.C. 112, paragraph 1, "Written Description" Requirement (66 FR 1099-1111, January 5, 2001) state,

"[p]ossession may be shown in a variety of ways including description of an actual reduction to practice, or by showing the invention was 'ready for patenting' such as by disclosure of drawings or structural chemical formulas that show that the invention was complete, or by describing distinguishing identifying characteristics sufficient to show that the applicant was in possession of the claimed invention" (*Id.* at 1104). Moreover, because the claims encompass a genus of variant species, an adequate written description of the claimed invention must include sufficient description of at least a representative number of species by actual reduction to practice, reduction to drawings, or by disclosure of relevant, identifying characteristics sufficient to show that Applicant was in possession of the claimed genus. However, factual evidence of an actual reduction to practice has not been disclosed by Applicant in the specification; nor has Applicant shown the invention was "ready for patenting" by disclosure of drawings or structural chemical formulas that show that the invention was complete; nor has Applicant described distinguishing identifying characteristics sufficient to show that Applicant were in possession of the claimed invention at the time the application was filed.

The *Guidelines* further state, "[f]or inventions in an unpredictable art, adequate written description of a genus which embraces widely variant species *cannot* be achieved by disclosing only one species within the genus" (Id. at 1106); accordingly, it follows that an adequate written description of a genus cannot be achieved in the absence of a disclosure of at least one species within the genus. As evidenced by Greenspan et al. (*Nature Biotechnology* 7: 936-937, 1999), defining epitopes is not as easy as it seems. Greenspan et al. recommends defining an epitope by the structural characterization of the molecular interface between the antigen and the antibody is necessary to define an "epitope" (page 937, column 2). According to Greenspan et al., an

epitope will include residues that make contacts with a ligand, here the antibody, but are energetically neutral, or even destabilizing to binding. Furthermore, an epitope will not include any residue not contacted by the antibody, even though substitution of such a residue might profoundly affect binding. Accordingly, it follows that the immunoepitopes that bind antibodies that can prevent biofilm formation can only be identified empirically. Therefore, absent a detailed and particular description of a representative number, or at least a substantial number of the members of the genus of immunoepitopes, the skilled artisan could not immediately recognize or distinguish members of the claimed genus antibodies that bind to the gacS protein and have the ability to prevent biofilm formation. Therefore, because the art is unpredictable, in accordance with the *Guidelines*, the description of immunoepitopes (antigenic determinants) is not deemed representative of the genus of antibodies to which the claims refer.

The rejection of claims 1-3 and 8 under 35 U.S.C. 112, first paragraph, as failing to comply with the enablement requirement is maintained for reasons of record. The claim(s) contains subject matter which was not described in the specification in such a way as to enable one skilled in the art to which it pertains, or with which it is most nearly connected, to make and/or use the invention.

Applicant argues:

- 1. The predictability in the art refers to the ability of the skilled artisan to extrapolate the disclosed results to the claimed invention.
- 2. One skilled in the art could readily anticipate the effect of the use of anti-gacS antibodies with the described knockout mutants of gacS disclosed in the specification and that antibodies of the

GacA/GacS regulatory system can be used to inhibit biofilm formation.

Applicant's arguments have been fully considered and deemed non-persuasive.

The rejection was made since the claimed invention was not described in the specification in such a way as to enable one skilled in the art to which it pertains, or with which it is most nearly connected, to make and/or use the invention without undue experimentation.

Undue experimentation is a conclusion reached by weighing the noted factual considerations set forth below as seen in *In re* Wands, 858 F.2d 731, 737, 8 USPQ2d 1400, 1404 (Fed. Cir. 1988). A conclusion of lack of enablement means that, based on the evidence regarding each of the factors below, the specification, at the time the application was filed, would not have taught one skilled in the art how to make and/or use the full scope of the claimed invention without undue experimentation.

The factors include, but are not limited to:

- 1. The breadth of the claims,
- 2. The nature of the invention,
- 3. The state of the prior art,
- 4. The level of one of ordinary skill,
- 5. The level of predictability in the art,
- 6. The amount of direction provided by the inventor,
- 7. The existence of working examples, and
- 8. The quantity of experimentation needed to make and/or use the invention based on the content of the disclosure.

Breadth of the claims

The rejected claims are drawn to the prophylactic use of antibodies that bind to the gacS polypeptide.

Application/Control Number: 10/828,557 Page 10

Art Unit: 1645

Working Examples/Guidance of Specification

The specification provides no working examples demonstrating the efficacy of claimed methods. The working examples are limited to methods of generating and evaluating gacS knockout mutants of *Pseudomonas chlororaphis* O6. The specification is silent with respect to the use of specific anti-gacS antibodies for the prevention of biofilm formation. Based on Applicant's assertion that one can predict the efficacy of a treatment regiment based on knockout mutants, one could prevent AIDS by the administration of anti-gp120 antibody since env-deficient HIV-1 mutants cannot effectively infect CD4+ T cells. However, to date, there is no AIDS "vaccine".

State of the prior art and Unpredictability of the art

The specification, as filed, is silent with regard to the claimed use of the claimed antibodies to prevent any type of biofilm. Applicant states that the claimed antibodies "... it is contemplated that inhibitors, antagonists or antibodies of the GacA/GacS regulatory system can also be used to inhibit biofilm formation..." in a prophetic sense but fails to demonstrate any prophylactic efficacy in any model system. The specification is silent as to which specific antibody, if any, would be effective to prevent the biofilm formation of a given microorganism. The examples, disclosed in the instant specification, are limited to methods of generating and evaluating gacS knockout mutants of *Pseudomonas chlororaphis* O6. While the skill in the art of immunology is high, to date, prediction of preventative efficacy for any given composition against any given microorganism is quite unpredictable. Given the lack of success in the art, the lack of working examples and the unpredictability of a given response (effect), the specification, as filed, does not provide enablement for methods of preventing biofilm formation, comprising the use of antibodies to a gacS protein.

The following is a quotation of the second paragraph of 35 U.S.C. 112:

Application/Control Number: 10/828,557

Art Unit: 1645

The specification shall conclude with one or more claims particularly pointing out and distinctly claiming the subject matter which the applicant regards as his invention.

The rejection of claims 1-3 and 8 under 35 U.S.C. 112, second paragraph, as being rendered vague and indefinite for failing to recite the active steps that need to be performed in the claimed method in order to achieve the stated goal of "preventing biofilm formation" is maintained for reasons of record. The amendment to claim 1 is insufficient to overcome this rejection as the claim still fails to recite the active steps that need to be performed in order to prevent biofilm formation.

Conclusion

No claim is allowed.

THIS ACTION IS MADE FINAL. Applicant is reminded of the extension of time policy as set forth in 37 CFR 1.136(a).

A shortened statutory period for reply to this final action is set to expire THREE MONTHS from the mailing date of this action. In the event a first reply is filed within TWO MONTHS of the mailing date of this final action and the advisory action is not mailed until after the end of the THREE-MONTH shortened statutory period, then the shortened statutory period will expire on the date the advisory action is mailed, and any extension fee pursuant to 37 CFR 1.136(a) will be calculated from the mailing date of the advisory action. In no event, however, will the statutory period for reply expire later than SIX MONTHS from the mailing date of this final action.

Application/Control Number: 10/828,557 Page 12

Art Unit: 1645

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Robert A. Zeman whose telephone number is (571) 272-0866. The examiner can normally be reached on Monday- Thursday, 7am -5:30 p.m.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Lynette Smith can be reached on (571) 272-0864. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free). If you would like assistance from a USPTO Customer Service Representative or access to the automated information system, call 800-786-9199 (IN USA OR CANADA) or 571-272-1000.

ROBERT A. ZEMAN PRIMARY EXAMINER

July 6, 2006